AREN34.US5.PCT PATENT

IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

In re Application: Robert M. Jones et al.

Confirmation No: 4098

Serial No.: 10/541,657

Group Art Unit: 1624

Filed: January 14, 2004(Intl. Filing Date)

Examiner: Not Yet Assigned

For: 1,2,3-TRISUBSTITUTED ARYL AND HETEROARYL DERIVATIVES AS MODULATORS OF METABOLISM AND THE PROPHYLAXIS AND TREATMENT OF DISORDERS RELATED THERETO SUCH AS DIABETES AND HYPERGLYCEMIA

Certificate of Mailing

I hereby certify that this correspondence, 22 citation sheets listing 386 references, and 342 references are being provided in seven (7) separate boxes, are being Deposited with the United States Postal Service as first class mail addressed to: Mail Stop PCT, Commissioner for Patents, Washington, DC 20231 On this date

February 27, 2007

By: A A A A B A B By: Susanne H. Goodson, Ph.D

Registration No.: 58,450

MAIL STOP PCT

Commissioner for Patents P.O. Box 1450 Alexandria, VA 22313-1450

Dear Sir:

INFORMATION DISCLOSURE STATEMENT

Pursuant to 37 C.F.R. §§ 1.56 and in accordance with 37 C.F.R. §§ 1.97 and 1.98, information relating to the above-identified application is hereby disclosed, the Examiner in charge of the above-identified application is requested to consider and make of record the references listed on the PTO Forms SB/08A and SB/08B, formerly known as PTO Form 1449, submitted herewith.

Inclusion of the information submitted herewith is not to be construed as an admission that the information is material as that term is defined in 37 C.F.R. § 1.56(b).

In accordance with 37 C.F.R. § 1.97(g), the filing of this Information Disclosure Statement shall not be construed to mean that a search has been made.

This 1	[nform	ation Disclosure Statement is being filed:
	within	n three months of the filing date of the patent application.
		three months of the date of entry into the national stage as set forth in F.R. § 1.491 of the international application.
\boxtimes	befor	e the mailing date of a first Office Action on the merits.
	after	the mailing date of a first Office Action on the merits, but before the
	mailir	ng date of a Final Office Action under 37 C.F.R. § 1.116 or a Notice of
	Allow	vance under 37 C.F.R. § 1.311, and accordingly is accompanied by:
		the Statement under 37 C.F.R. § 1.97(e) (see "Statement" below);
		or
		the Fee of \$180.00 set forth in 37 C.F.R. § 1.17(p); or
		No fee is owed by the applicant(s).
	In acc	ordance with 37 C.F.R. § 1.129(a), this Information Disclosure
	Stater	nent is being filed in connection with the first or second After
	Final	Submission, and accordingly is accompanied by the Statement under 37
	C.F.R	§ 1.97(e) (see "Statement" below) and the fee of \$180.00 as set forth in
	37 C.	F.R. § 1.17(p), is attached.
	after	the mailing date of a Final Office Action under 37 C.F.R. § 1.116 or a
	Notic	e of Allowance under 37 C.F.R. § 1.311, but before the payment of the
	Issue	Fee, and accordingly is accompanied by the Statement under 37 C.F.R.
	§ 1.97	(e), (see "Statement," and "Fees" below).
\boxtimes	Copie	es of the references (excluding the U.S. Patent Documents) listed on the
	attach	ed PTO Forms SB/08a and SB/08b, formerly known as PTO Form 1449
	are en	closed.
	EXC	EPT THAT:
	\boxtimes	In view of the voluminous nature of references JF, LX, ME, PI, PJ,
		and, PK and the likelihood that these references are available to the
		Examiner, copies are not enclosed herewith.

AREN34.US5.PCT

PATENT

No fee or Statement is required under 37 C.F.R. § 1.97(b) as no first Office Action on the merits has been received by Applicants.

Respectfully submitted,

Susanne Hoff Goodson, Ph.D.

Registration No. 58,450

Dated: February 27, 2007

COZEN O'CONNOR, P.C. 1900 Market Street, 5th Floor Philadelphia, PA 19103-3508 (215) 665-2000 - Telephone (215) 665-2013 - Facsimile U.S. Patent and Trademark Office; U.S. DEPARTMENT OF COMMERCE Under the Paperwork Reduction Act of 1995, no persons are required to respond to a collection of information unless it contains a valid OMB control number.

Substitute for form 1449A/PTO

Sheet

INFORMATION DISCLOSURE STATEMENT BY APPLICANT

(Use as many sheets as necessary)

of

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Application Number 10/541,657

Filing Date January 14, 2004

First Named Inventor Robert M. Jones

Art Unit 1624

Examiner Name To Be Determined

Attorney Docket Number AREN34.US5.PCT

			U.S. PATENT D	OCUMENTS	
Examiner Initials *	Cite No. ¹	Document Number Number - Kind Code ² (if known)	Publication/Issue Date MM-DD-YYYY	Name of Patentee or Applicant of Cited Document	Pages, Columns, Lines, Where Relevant Passages or Relevant
	AA	US-3,503,963	03-31-1970	Schweizer, et al.	Figures Appear
	AB	US-3,592,932	07-13-1997	Duerr et al.	
<u> </u>	AC	US-3,608,087	09-21-1971	Patchett et al.	
	AD	US-3,686,238	08-22-1972	Zaffaroni et al.	
 ,	AE	US-3,690,834	09-12-1972	Goldstein et al.	
<u> </u>	AF	US-3,849,420	11-19-1974	Tong	
	AG	US-3,852,434	12-03-1974	Kahan et al.	
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_	AI	US-3,887,329	06-03-1975	Hegar et al.	
•	AJ	US-3,966,744	06-29-1976	Goldstein et al.	
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- · · · · ·	AN	US-4,101,541	07-18-1978	Petitpierre et al.	
-	AO	US-4,189,427	02-19-1980	Komorowski	
-	AP	US-4,242,507	12-30-1980	Itoh et al.	
	AQ	US-4,267,174	05-12-1981	Berger et al.	
	AR	US-4,275,148	06-23-1981	Endo et al.	
	AS	US-4,397,848	08-09-1983	Bosies et al.	
	AT	US-4,517,183	05-14-1985	Bosies et al.	
	AU	US-5,691,364	11-25-1997	Buckman et al.	
	AV	US-5,849,759	12-15-1998	Arnaiz et al.	
	AW	US-5,948,786	09-07-1999	Fujiwara et al.	

FOREIGN PATENT DOCUMENTS

		Foreign Patent Document			Pages, Columns, Lines,	
Examiner Initials*	Cite No. ¹	Country Code ³ - Number ⁴ - Kind Code ⁵ (<i>if known</i>)	Publication Date/Filing Date MM-DD-YYYY	Name of Patentee or Applicant of Cited Document	Where Relevant Passages or Relevant Figures Appear	Τ ⁶
	AX	AU 492126	11-20-1975	Ciba-Geigy AG		
	AY	AT 327605 (w/Eng. abst)	06-15-2006	Deutsche Telekom AG		
	AZ	BE 829845 (w/counterpart USP 3,984,411)	12-04-1975	Societe Generale de Recherches et d'applications scientifiques		
	BA	BE 868796 (w/counterpart USP 4,267,174)	01-08-1979	Boehringer		<u></u>
	BB	CH 560197 (w/Eng. abst.)	03-27-1975	Ciba-Geigy AG		
	BC	DE 19602095 (w/Eng. abst)	07-24-1997	Bayer AG (DE)		
	BD	DE 19737723 (w/Eng. abst)	02-18-1999	Bayer AG (DE)		
	BE	DE 19962936 (w/Eng. abst)	06-28-2001	Bayer AG (DE		

Examiner Signature Date Considered

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Complete if Known Substitute for form 1449A/PTO 10/541,657 **Application Number** INFORMATION DISCLOSURE January 14, 2004 Filing Date STATEMENT BY APPLICANT Robert M. Jones First Named Inventor 1624 Art Unit To Be Determined (Use as many sheets as necessary) Examiner Name AREN34.US5.PCT Sheet of 22 Attorney Docket Number

			U.S. PATENT D	OCUMENTS	
Examiner	Cite	Document Number	Publication/Issue Date	Name of Patentee or Applicant of Cited Document	Pages, Columns, Lines, Where Relevan
Initials *	No. ¹	Number - Kind Code ² (if known)	MM-DD-YYYY		Passages or Relevant Figures Appear
	BF	US-5,962,479	10-05-1999	Chen	
- · · -	BG	US-6,008,234	12-28-1999	Kochanny et al.	
	BH	US-6,187,777	02-13-2001	Norman et al.	
	BI	US-6,218,431	04-17-2001	Schoen et al.	
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	BK	US-6,414,002	07-02-2002	Cheng et al.	
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,	ВМ	US-6,545,016	04-08-2003	Dellaria et al.	
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····	ВО	US-6,583,154	06-24-2003	Norman et al.	
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	BQ	US-6,956,047	10-18-2005	Chen et al.	
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		FOREIGN	PATENT DOC	UMENTS		
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Examiner Initials*	Cite No. ¹	Country Code ³ - Number ⁴ - Kind Code ⁵ (<i>if known</i>)	Publication Date/Filing Date MM-DD-YYYY	Name of Patentee or Applicant of Cited Document	Relevant Passages or Relevant Figures Appear	T ⁶
	BT	DE 2048375 (w/GB1311956)	04-22-1971	Merck & Co.		
	BU	DE 2223644 (w/GB1393993)	11-30-1972	Ciba-Geigy AG		
	BV	DE 2341925 (w/Eng. abst)	03-06-1975	Thomae Gmbh		
<u> </u>	BW	DE 2356644 (w/USP3,948,914)	05-22-1974	Ciba Geigy AG		
	BX	DE 2460238 (w/GB1493380)	07-03-1975	Ciba Geigy AG		
	BY	DE 2503136 (w/GB1495665)	07-31-1975	Products Chimiques Ugine Kuhlmann		
	BZ	DE 2831850 (w/USP 4,273,870)	02-07-1980	BASF AG		
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	СВ	DE 3406329 (w/Eng. abst.)	08-22-1985	Merck & Co.		

Examiner	Date	
Signature	Considered	
3		

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Sheet	3	of	22	Attorney	Docket Number	AREN34.U	S5.PCT	
			FC	DREIGN PAT	ENT DOCUME	NTS		
Examin er Initials*	Cite No. ¹	Foreign Pat Country Code ³ - Num <i>known</i>)	ent Document ober ⁴ - Kind Code ⁵ (<i>if</i>	Publication Date/Filing Date MM-DD-YYYY		or Applicant of Cited ument	Pages, Columns, Lines, Where Relevant Passages or Relevant Figures Appear	Τ ⁶
	CC	DE 3601196 (v	v/USP4,766,213)	07-23-1987	Mercl	k & Co.		
	CD	EP 0 014 976 (w/USP4,517,183)	09-03-1980	Boeh	ringer		
	CE	EP 0 055 693 (v	w/USP4,493,726)	07-07-1982	CIBA C	Geigy AG		
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	CJ	EP 0 565 488	(w/Eng. abst.)	10-13-1993	Ciba G	eigy AG		
	CK	EP 0 604 800	(w/Eng. abst.)	07-06-1994	Thoma	e BMGH		
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	CP	EP 1 (074 549	02-07-2001	F. Hoffmann	La Roche AG		<u> </u>
	CQ	EP 1	040 831	05-23-2003	Pf	izer		
	CR	EP 0 149 088 (v	w/USP4,643,995)	12-01-1984	Degus	ssa Akt.		
	CS	EP 0	191 603	08-20-1986	Fujisawa	a Pharma		<u>.</u>
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	CU	EP 1	340 749	09-03-2003	Takada Chem	nical Industries		
	CV	EP 1	475 094	11-10-2004	Ustav Ex Bo	tan Akademie		ļ <u> </u>
	CW	FR 1551400 (w	v/USP3,598,801)	12-27-1968	J.R. Ge	eigy AG		
	CX	GB 9	35595	08-28-1963	Cib	a Ltd		
	CY	GB 1	311956	03-28-1973	Mercl	с & Co.		
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	DC		w/2 Eng. absts.)	02-06-1980		F AG		
	DD		0 (w/Eng. abst.)	02-08-2000	<u> </u>	harmaceutical		
	DE		2 (w/Eng. abst.)	04-04-2001		уо Со		<u> </u>
	DF		w/Eng. translation)	09-30-2004		Pharmaceutical		<u> </u>
	DG		w/Eng. translation	09-30-2004		Pharmaceutical		<u> </u>
	DH		v/USP3,503,963)	04-24-1967		imited		<u> </u>
	DI	NL 6814810 (w/GB1250624)	04-21-1969		Limited		<u> </u>
	DJ	SU 938 559	(w/Eng. abst.	11-30-1993		nyj Ni Khim natsevt		
	DK	WO 9	4/13677	06-23-1994	Pf	izer		
Examine Signature					Date Considered			

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Substitut	e for form 1449	PA/PTO .		Complete if Known			
				Application Number	10/541,657		
INFO	PRMATI	ON DIS	CLOSURE	Filing Date	January 14, 2004		
STA	TEMEN'	T BY A	PPLICANT	First Named Inventor	Robert M. Jones		
				Art Unit	1624		
	(Use as ma	ny sheets as	necessary)	Examiner Name	To Be Determined		
Sheet	4	of	22	Attorney Docket Number	AREN34.US5.PCT		

Sneet	4	FOREI	GN PATENT I	DOCKEL NUMBER	AREN34.030		
			GNPATENTI	DOCUMENTS		Pages, Columns, Lines,	<u> </u>
Examiner Initials*	Cite No. ¹	Foreign Patent Document Country Code ³ - Number ⁴ - Kind Code ⁵ (if known)	Publication Date/Filing Date MM-DD-YYYY		ntee or Applicant of Cited Document	Where Relevant Passages or Relevant Figures Appear	Т
	DL	WO 95/33750	12-14-1995		Pfizer		
	DM	WO 96/28427	09-19-1996	В	erlex Lab		
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	DR	WO 97/26252	07-24-1997	FI	MC Corp.		
	DS	WO 97/29109	08-14-1997	Janssen Ph	armaceutica, et al.		
	DT	WO 97/49706	12-31-1997	No	vartis AG		
	DU	WO 98/04528	02-05-1998	Ba	yer Corp.		
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	EF	WO 00/35875	06-22-2000	Am H	ome Products		1
	EG	WO 00/35886	06-22-2000	Axy	s Pharm Inc		
	EH	WO 01/22938	04-05-2001	Janssen	Pharmaceutica	- 1 11111111111111111111111111111111111	
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	EL	WO 01/27107	04-19-2001		Myers Squibb		1
	EM	WO 01/47887	07-05-2001		ayer Akt.	<u> </u>	1
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	EO	WO 01/53263	2001-07-26		er Products		1
	EP	WO 01/58900 (w/Eng. abst.)	08-16-2001		riku Seiyaku		
Examiner Signature				Date Considered			

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Substitute for form 1449A/PTO Complete if Known 10/541,657 Application Number INFORMATION DISCLOSURE January 14, 2004 Filing Date STATEMENT BY APPLICANT Robert M. Jones First Named Inventor 1624 Art Unit To Be Determined (Use as many sheets as necessary) Examiner Name AREN34.US5.PCT 22 Sheet 5 Attorney Docket Number of

	γ		N PATENT DO	COMEN 19	Constant Contract Constant	7
Examiner Initials*	Cite No. ¹	Foreign Patent Document Country Code ³ - Number ⁴ - Kind Code ⁵ (if known)	Publication Date/Filing Date MM-DD-YYYY	Name of Patentee or Applicant of Cited Document	Pages, Columns, Lines, Where Relevant Passages or Relevant Figures Appear	Т
	EQ	WO 01/62233	08-30-2001	F. Hoffmann La Roche		
	ER	WO 01/85699	11-15-2001	Janssen Pharmaceutica		
	ES	WO 02/02549	01-10-2002	Taisho Pharma		
-	ET	WO 02/06237 (w/Eng. abst.)	01-24-2002	Yamanouchi Pharma		
	EU	WO 02/06274	01-24-2002	American Home Prod		
	EV	WO 02/070485 (w/Eng. abst.)	09-12-2002	Bayer Akt.		
	EW	WO 02/072101	09-19-2002	Bristol-Myers Squibb		
	EX	WO 02/19975 (w/Eng abst.)	03-14-2002	Taisho Pharma.	<u>.</u>	
	EY	WO 02/32893	04-25-2002	Schering Corp.		
	EZ	WO 02/40451	05-23-2002	Eli Lilly & Co.		
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	FD	WO 02/44362 (w/Eng Abst)	06-06-2002	Yamanouchi Pharma		
	FE	WO 02/59083	08-01-2002	Smithkline Beecham		
•	FF	WO 02/98864	12-12-2002	F. Hoffmann La Roche		
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	FJ	WO 03/026661 (w/Eng. abst.)	04-03-2003	Yamanouchi Pharma		
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	FQ	WO 2004/000819	12-31-2003	AstraZeneca AB		
	FR	WO 2004/000843	12-31-2003	AstraZeneca AB		_
	FS	WO 2004/009596	01-29-2004	SmithKline Beecham		<u> </u>
	FT	WO 2004/009597	01-29-2004	SmithKline Beecham		<u> </u>
	FU	WO 2004/009602	01-29-2004	SmithKline Beecham		<u> </u>
·	FV	WO 2004/024943 (w/Eng. Abst.)	03-25-2004	Yamanouchi Pharma		<u> </u>
	FW	WO 2004/029204	04-08-2004	Merck & Co.		<u> </u>
	FX	WO 2004/031189	04-15-2004	Bristol Myers Squibb		
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Substitu	te for form 1449A/PT0	0			Complete if Known
				Application Number	10/541,657
INF	DRMATION	DIS	CLOSURE	Filing Date	January 14, 2004
STA	STATEMENT BY APPLICANT		PPLICANT	First Named Inventor	Robert M. Jones
				Art Unit	1624
	(Use as many she	eets as	necessary)	Examiner Name	To Be Determined
Sheet	6	of	22	Attorney Docket Number	AREN34.US5.PCT

		FOREIG	N PATENT D	OCUMENTS		_
Examiner Cite No.1		Foreign Patent Document Country Code ³ - Number ⁴ - Kind Code ⁵ (if known)	Publication Date/Filing Date MM-DD-YYYY	Name of Patentee or Applicant of Cited Document	Pages, Columns, Lines, Where Relevant Passages or Relevant Figures Appear	۲°
	FY	WO 2004/035588	04-29-2004	SmithKline Beecham		
•	FZ	WO 2004/041164	05-21-2004	Merck & Co.		
	GA	WO 2004/056825	07-08-2004	Syngenta		
	GB	WO 2004/056829	07-08-2004	Syngenta		_
	GC	WO 2004/062665	07-29-2004	SB Pharmco et al.		
	GD	WO 2004/065380	08-05-2004	Arena Pharm.		
•	GE	WO 2004/074218	09-02-2004	Avanir Pharmaceuticals		
	GF	WO 2004/076413	09-10-2004	Arena Pharm.	,	-
***	GG	WO 2004/111000	12-23-2004	Fujisawa Pharmaceutical		
٠	GH	WO 2005/016894	02-24-2005	Novartis Ag		
	GI	WO 2005/030129	04-07-2005	Merck & Co.		
	GJ	WO 2005/035525	04-21-2005	Vertex Pharm		
	GK	WO 2005/037215	04-28-2005	Massachusetts Inst of Technology		
	GL	WO 2005/046603	05-26-2005	Synta Pharmaceuticals		
· · · · · · · · · · · · · · · · · · ·	GM	WO 2005/049033	06-02-2005	AstraZeneca AB		
	GN	WO 2005/058315	06-30-2005	Ribapharm Inc		
	GO	WO 2005/061489	07-07-2005	Prosidion Ltd.		
·· · · · ·	GP	WO 2005/090348	09-29-2005	Glaxo Group Ltd		•••
	GQ	WO 2005/100365 (w/Eng. abst.)	10-27-2005	Sankyo Co Ltd.		
	GR	WO 2005/117909	12-15-2005	Exelixis Inc		
	GS	WO 2006/067531	06-29-2006	Prosidion Ltd.		
	GT	WO 2006/067532	06-29-2006	Prosidion Ltd.		
	GU	WO 2006/040966	04-20-2006	Astellas Pharma		
	GV	WO 2006/043490 (w/Eng. Abst)	04-27-2006	Astellas Pharma		
	GW	WO 2006/070208	07-06-2006	Prosidion Ltd.		

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		FOREIG	N PATENT D	OCUMENTS		
Examiner Initials*	4	Foreign Patent Document Country Code ³ - Number ⁴ - Kind Code ⁵ (if known)	Publication Date/Filing Date MM-DD-YYYY	Name of Patentee or Applicant of Cited Document	Pages, Columns, Lines, Where Relevant Passages or Relevant Figures Appear	Τ ⁶
	00	WO 2005/075426	08-18-2005	Glenmark Pharm		
	OP	WO 2005/072530	08-11-2005	Merck		
	OQ	WO 2005/063750	07-14-2005	Boehringer		
	OR	WO 2005/058849	06-30-2005	Glenmark Pharm		
	OS	WO 2005/047297	05-26-2005	Phenomix Corp.		
···	OT	WO 2005/042488	05-12-2005	Takeda Pharm		· ·
	OU	WO 2005/040095	05-06-2005	Astrazeneca		
	OV	WO 2005/033099	04-14-2005	Glenmark Pharm		
	OW	WO 2005/030751	04-07-2005	Syrrx		
	OX	WO 2005/030127	04-07-2005	Merck		
	OY	WO 2005/026148	03-24-2005	Syrrx		
	OZ	WO 2005/025554	03-24-2005	Japan Tobacco		
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•	PF	WO 97/40832	11-06-1997	Hans-Knoll-Inst		
· · · ·	PG	WO 2005/121121	12-22-2005	Arena Pharm		
•	PH	WO 2005/007647	01-27-2005	Arena Pharm		

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Substitut	e for form 1449E	В/РТО		Complete if Known				
W156		D.O		Application Number	10/541,657			
		_	CLOSURE	Filing Date	January 14, 2004			
STA	TEMENT	BYA	PPLICANT	First Named Inventor	Robert M. Jones			
				Art Unit	1624			
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· ·		NON PATENT LITERATURE DOCUMENTS						
Examiner Initials *	Cite No. Include name of the author (in CAPITAL LETTERS), title of the article (when appropriate), title of the item (book, magazine, journal, serial, symposium, catalog, etc.), date, page(s), volume-issue number(s), publisher, city and/or country where published.							
	GX	ABDALLA et al., "Synthesis and reaction of 3-cyano 2-(1H)-pyridones," Pakistan Journal of Scientific and Industrial Research (1977) 20(3):139-149.						
	GY	ABRAMOVITCH et al., "Solution and flash vacuum pyrolysis of some 2,6-disubstituted β -phenethylsulfonyl azides and of β -styrenesulfonyl azide," J Org Chem (1985) 50:2066-2073.						
	GZ	APPUKKUTTAN et al., "Transition-Metal-Free Sonogashira-Type Coupling Reactions In Water," European Journal Of Organic Chemistry (2003) 24:4713-4716.						
	НА	ARVANITIS et al., "Non-peptide corticotropin-releasing hormone antagonists: syntheses and structure-activity relationships of 2-anilinopyrimidines and -triazines.," J Med Chem. (1999) 42(5):805-18.						
	НВ	ARVANITIS et al., "Non-peptide corticotropin-releasing hormone antagonists: syntheses and structure-activity relationships of 2-anilinopyrimidines and -triazines.," J Med Chem. (1999) Supporting Material, pp. 1-10.						
	НС	ARVANITIS et al., "CRF Ligands via suzuki and negishi couplings of 3-pyridyl boronic acids or halides with 2-benzyloxy-4-chloro-3-nitropyridine," <i>Bioorganic & Medicinal Chemistry Letters</i> (2003) 13(2):289-291.	_					
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	нн	BAKKESTUEN et al., "Regioselective N-9 arylation of purines employing arylboronic acids in the presence of Cu(II)," Tetrahedron Letters (2003) 44:3359-3362.						
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Examiner Signature		Date Considered						

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Applicant's unique citation designation number (optional). ² Applicant is to place a check mark here if English language Translation is attached. This collection of information is required by 37 CFR 1.98. The information is required to obtain or retain a benefit by the public which is to file (and by the USPTO to process) an application. Confidentiality is governed by 35 U.S.C. 122 and 37 CFR 1.14. This collection is estimated to take 120 minutes to complete, including gathering, preparing, and submitting the completed application form to the USPTO. Time will vary depending upon the individual case. Any comments on the amount of time you require to complete this form and/or suggestions for reducing this burden, should be sent to the Chief Information Officer, U.S. Patent and Trademark Office, U.S. Department of Commerce, P.O. Box 1450, Alexandria, VA 22313-1450. DO NOT SEND FEES OR COMPLETED FORMS TO THIS ADDRESS. SEND TO: Commissioner for Patents, P.O. Box 1450, Alexandria, VA 22313-1450.

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communication to applicant.

Applicant's unique citation designation number (optional). Applicant is to place a check mark here if English language Translation is attached.

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Substitute		149B/PTO	T	Complete if Known							
			Application Number	10/541,657							
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STAT	EME	NT BY APPLICANT	First Named Inventor Robert M. Jones								
			Art Unit	1624							
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		NON PATEN	T LITERATURE DOCUMENTS								
Examiner Initials *	Cite No.1	Include name of the author (in CAPITAL LETTERS), title of the article (when appropriate), title of the item (book, magazine, journal, serial, symposium, catalog, etc.), date, page(s), volume-issue									
	IA		BULGER et al., "An investigation into the alkylation of 1,2,4-triazole," Tetrahedron Letters (2000)								
	IB	CHAN et al., "Isoquinoline-6-Carbo (HCMV)Inhibitors," Bioorganic & M									
	IC	CHEN et. al., "Optimization of 3-ph releasing factor-1 antagonists with a Medicinal Chemistry Letters (2004) 14	adequate lipophilicity and wate	nidines as potent corticotrophiner solubility," <i>Bioorganic &</i>							
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	IG	CLARK et al., "Synthesis and Analg b]pyridine-2-ones and 3-(Substitute 21(9):965-978.	gesic Activity of 1,3-Dihydro-3-d phenyl)-1,2,3-triazolo(4,5-b]p	(Substituted phenyl)imidazo[4,5-yridines," J. Med. Chem. (1978)							
	IH	COCUZZA et al., "Use of the Suzul Corticotropin-Releasing Hormone ((1999) 9:1063-1066.									
	П	COHEN et al., "The Preparation an Chemistry, Sloan-Kettering Institute for Sci., Cornell Uiv. Med. College (1962)	or Cancer Research, and Sloan Kel								
	IJ	COLANDREA et al., "Synthesis and Tetrahedron Letters (2000) 41:8053-80	•	6- and 1,7-naphythridines,"							
	IK	COLLIER et al., "Radiosynthesis an [125]]-ITIPP(Ψ)," J. Labeled Compd. Re									
	ΠL	COSSEY et al., "Amide-acid chlorid cyanoacetamides," Australian Journal									
	IM	GS39783 (N,N'-dicyclopentyl-2-met activity without side effects associa and Experimental Therapeutics (2004)	terization of the novel GABAB receptor-positive modulator ethylsulfanyl-5-nitropyrimidine-4,6-diamine): Anxiolytic-like ated with baclofen or benzodiazepines," Journal of Pharmacology								
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		NON PATENT LITERATURE DOCUMENTS							
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	Ю	DESIMONI et al., "Polynuclear Isoxazole Types-I – Isoxazolo[4,5-d]Pyrimidines," Tetrahedron (1967) 23:675-680.							
	IP	DEVITA et al., "Identification and initial structure-activity relationships of a novel non-peptide quinolone GnRH receptor antagonist," Bioorg & Med Chem Ltrs (1999) 9(17):2615-2620.							
	IQ	DI BRACCIO et al., "Synthesis and preliminary pharmacological examination of 2,4-disubstituted N,N-dialkyl-1,8-naphthyridine-3-carboxamides," Farmaco (1989) 44(9):865-881.							
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	IS	EICHER et al., "Reaction of triafulvenes with isonitriles. A simple synthesis of diphenyl-substituted functionalized cyclobutene derivatives and related products," Synthesis (1987) (7):619-626.							
	ESCHER et al., "Cyclopentylamine Substituted Triazolo[4,5-D]Pyrimidine: Implications for Binding to the Adenosine Receptor," <i>Tetrahedron Letters</i> (1991) 32(29):3583-3584.								
	IV	GANGLOFF et al., "Synthesis of 3,5-disubstituted-1,2,4-oxadiazoles using tetrabutylammonium fluoride as a mild and efficient catalyst," <i>Tetrahedron Letters</i> (2001) 42:1441-1443.							
	IW	GILLIGAN et al., "Corticotropin-releasing factor antagonists: Recent advances and exciting prospects for the treatment of human diseases," Current Opinion in Drug Discovery & Development (2004) 7(4):487-497.							
	IX	GILLIGAN, et al., "Corticotropin Releasing Factor (CRF) Receptor Modulators" Progress and Opportunities for New Therapeutic Agents," J. Med. Chem. (2000) 43(9):1641-1660.							
	ſΥ	GOLDNER et al., "Die Darstellung 2,9-; 2,6,9- und 6,9-substituierter Purine," Journal fuer Praktische Chemie (Leipzig) (1961) 12:242-252.							
	IZ	GINER-SOROLLA et al., "The Synthesis and Properties of 6-Mercaptomethylpurine and Derivatives," Cornell University Medical College (1965) 8:667-672.							
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	HAMADA et al., "An improved synthesis of arylsulfonyl chlorides from arylhalides," Synthesis (1986) pp. 852-854.								
	JC	HE et al., "4-(1,3-Dimethozyprop-2-ylamino)-2,7-dimethyl-8-(2,4-dichlorophenyl)-pyrazolo[1,5-a]- 1,3,5-triazine: A Potent, Orally Bioavailable CRF1 Receptor Antagonist," J. Med. Chem. (2000) 43:449- 456.							
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Complete if Known Substitute for form 1449B/PTO 10/541,657 **Application Number** INFORMATION DISCLOSURE January 14, 2004 Filing Date STATEMENT BY APPLICANT First Named Inventor Robert M. Jones Art Unit 1624 To Be Determined (Use as many sheets as necessary) Examiner Name AREN34.US5.PCT 22 12 Attorney Docket Number of Sheet NON PATENT LITERATURE DOCUMENTS Include name of the author (in CAPITAL LETTERS), title of the article (when appropriate), title of T² the item (book, magazine, journal, serial, symposium, catalog, etc.), date, page(s), volume-issue Cite Examiner number(s), publisher, city and/or country where published. No. Initials * HECHT et al., "On the "activation" of cytokins*," J of Biological Chemistry (1975) 250(18):7343-7351. JD HERSPERGER et al., "Palladium-Catalyzed Cross-Coupling Rtions for the Synthesis of 6,8-Disubstituted 1,7-Naphthyridines: A Novel Class of Potent and Selective Phosphodiesterase Type JE 4D Inhibitors," J. Med. Chem. (2000) 43:675-682. HIGUCHI et al., "Pro-drugs as novel delivery systems," A.C.S. Symposium Series, Vol. 14 (1987). *JF HILL et al., "Environmental contributions to the obesity epidemic," Science (1998) 280(5368):1371-4. JG HOCEK et al., "An Efficient Synthesis of 2-Substituted 6-Methylpurine Bases and Nucleosides by Fe- or Pd-Catalyzed C ross-Coupling Reactions of 2,6-Dichloropurines," J. Org. Chem. (2003) JH 68:5773-5776. HUANG et al., "Synthesis and Antiplatelet Activity of Phenyl Quinolones," Bioorganic & Medicinal Л Chemistry (1998) 6:1657-1662. BERGE et al., "Pharmaceutical Salts," Journal of Pharmaceutical Sciences (1977) 66(1):1-19. IJ JIA, et al., "Design, Synthesis and Biological Activity of Novel Non-Amidine Factor Xa Inhibitors. Part 1: P1 Structure-Activity Relationships of the Substituted 1-(2-Naphtyl)-1H-pyrazole-5-JK carboxylamides," Bioorganic & Medicinal Chemistry Letters (2002) 12:1651-1655. JOGIE et al., "Unusual protein-binding specificity and capacity of aza-arenophilic gels," Journal of JL Molecular Recognition (1998) 11:261-262. KAWASE et al., " α -trifluoromethylated acyloins induce apoptosis in human oral tumor cell lines," JM Bioorg & Med Chem Ltrs (1999) 9(21):3113-3118.

Signature *EXAMINER: Initial if reference considered, whether or not citation is in conformance with MPEP 609. Draw line through citation if not in conformance and not considered. Include copy of this form with next communication to applicant.

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KELLY et al., "A Synthesis of Aaptamine," Tetrahedron (1985) 41(15):3033-3066.

6-(dimethylamino)-9H-purines," J Med Chem (1990) 33(1):196-202.

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Substitute	for form 1449	В/РТО			Complete if Known			
INICO				Application Number	10/541,657			
			CLOSURE	Filing Date	January 14, 2004			
STAT	LEMEN.	T BY A	PPLICANT	First Named Inventor	Robert M. Jones			
				Art Unit	1624			
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	JР				• •		rs of phosphodiestera inal Chemistry Letters		PDE7): synthesis and initial 15:1829-1833.	
	JQ			_			•	•	on 2 and 4. Nucleophilic emistry (1994) 131(3-4):521-527.	
	JR	KLOET 96(5):15			orierendė formy	/lierungsreak	tionen an pyrimidine	en," M	onatshefte fuer Chemie, (1965)	
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	JU	1	KRAUZE et al., "Synthesis of 3-oxoisothiazolo[5,4-b]pyridines," Khimiya Geterotsiklicheskikh Soedinenii (1982) (4):508-512.							
	JV	tetrahy	KUMEGAI et al., "Synthesis, SAR and biological activities of CRH1 Receptor: Novel 3- or 4-carbamoyl-1,2,5,6-tetrahydropyridinopyrrolopyrimidine derivative," 4th ACS National Meeting, August 18-22, 2002, Boston, MA. Poster #259.							
	JW		•	•	nethod for the el mmunications (20		•	yl-subs	stituted methyl alcohols into	
	JX	LANII 12(11):			ll molecule co	rticotrophi	n-releasing factor a	antago	onists," Expert Opinion (2002)	
	JΥ	LEADBEATER et al., "First Examples Of Transition-Metal Free Sonogashira-Type Couplings," Organic Letters (2003) 5(21):3919-3922								
Examiner Signature			Date Considered							

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			CLOSURE	Filing Date	January 14, 2004			
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				Art Unit	1624			
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	JZ						onogashira-type co ation, pp. S1-S4.	ouplings,"	Department of Chemistry,	_
	KA		E et al., "Synthesis and biological evaluation of clitocine analogues as adenosine kinase hibitors," Bioorg & Med Chem Ltrs (2001) 11(18):2419-2422.							
	KB				ial antipurines. I Society (1958)	•		stituted pu	irines and 8-azapurines,"	
	KC						00652218 for the exern. (2001) 44:S280-	_	of the tachykinin NK1	
	KD				arly changes in obesity," Diabe	-	ndial insulin secreti) 43:696-702.	on, not in i	insulin sensitivity,	
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	KI	LUO 6 5742.	et al., "	'Microwa	eve-assisted syr	ithesis of	aminopyrimidines	" Tetrahedi	ron Letters (2002) 43:5739-	
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	KK			Mild Met) 5(14):24		n Coupli	ng Reaction of Am	nes and A	ryl Halides," Organic	
	KL				·		ed 3-aryl- and 3-cy Chem (2003) 46(1):		peridines as partial	
	MACKMAN et al., "2-(2-Hydroxy-3-alkoxyphenyl)-1H-benzimidazole-5-carboxamidine derivati as potent and selective urokinase-type plasminogen activator inhibitors," Bioorganic & Medicinal Chemistry Letters (2002) 12(15):2019-2022.									
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		NON PATENT LITERATURE DOCUMENTS	
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	KN	MAJEED, et al, "Stannylation Reactions and Cross-Couplings in Pyrimidines," Tetrahedron (1989) 45(4):993-1006.	
	ко	MATSUI et al., "Highly potent inhibitors of TNF- α production. Part II: metabolic stabilization of a newly found chemical lead and conformational analysis of an active diastereoisomer," <i>Bioorg Med Chem.</i> (2002) 10(12):3787-805.	
	KP	MATSUNO et al., "Potent and selective inhibitors of platelet-derived growth factor receptor phosphorylation. 3. Replacement of quinazoline moiety and improvement of metabolic polymorphism of 4-[4-(N-substituted (thio)carbamoyl)-1-piperazinyl]-6,7-dimethoxyquinazoline derivatives," J Med Chem (2003) 46(23):4910-4925.	
	KQ	MESGUICHE et al., "4-Alkoxy-2,6-diaminopyrimidine derivatives: inhibitors of cyclin dependent kinases 1 and 2," Bioorganic & Medicinal Chemistry Letters (2003) 13(2):217-222.	
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	KS	MITTELBACH et al., "Syntheses with nitriles. 60. Preparation of 4-amino-5-cyano-6-phenylpyrimidines from 2-amino-1,1-dicyano-2-phenylethene," Journal of Heterocyclic Chemistry (1980) 17(7):1385-1387.	
	KT	MIYASHITA et al., "Preparation of Heterarenecarbonitriles by Reaction of Haloheteroarenes with Potassium Cyanide Atalyzied by Sodium p -Toluenesulfinate," Heterocycles (1994) 39(1):345-350.	
	KU	MOHAN et al., "Solid-phase synthesis of N-substituted amidinophenoxy pyridines as factor Xa inhibitors," Bioorganic & Medicinal Chemistry Letters (1998) 8(14):1877-1882.	
	KV	MOMBEREAU et al., "Genetic and Pharmacological Evidence of a Role for GABAB Receptors in the Modulation of Anxiety- and Antidepressant-Like Behavior," <i>Neuropsychopharmacology</i> (2004) 29(6):1050-1062.	
	KW	MONGIN et al., "Advances in the directed metallation of azines and diazines (pyridines, pyrimidines, pyridazines, quinolines, benzodiazines and carbolines). Part 1: Metallation of pyridines, quinolines and carbolines," <i>Tetrahedron</i> (2001) 57(19):4059-4090.	
	KX	MONTGOMERY et al., "Isonucleosides. I. Preparation of methyl 2-deoxy-2-(purin-9-yl)arabinofuranosides and methyl 3-deoxy-3-(purin-9-yl)xylofuranosides," Journal of Organic Chemistry (1975) 40(13):1923-1927.	
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STAT	TEMENT	BY A	PPLICANT	First Named Inventor	Robert M. Jones		
				Art Unit	1624		
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Sheet	16	of	22	Attorney Docket Number	AREN34.US5.PCT		

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	KZ	MOSCHITSKII et al., Translation of "Reaction of 2,3,5,6-tetrachloro-4-pyridyl-vinyl sulfone with nuleophilic agents," <i>Khimiya Geterotsiklicheskikh Soedinenii</i> (1972) pp. 1634-1637, (Translated Pages 1482-1485).					
	LA	MULLER et al., "7-Deaza-2-phenyladenines: Structure-Activity Relationships of Potent A1 Selective adenosine Receptor Antagonists," J. Med. Chem. (1990) 33:2822-2828					
	LB	NAKAZATO et al., "Synthesis, SAR and biological activities of CRH1 Receptor: Novel 3- or 4-carbamoyl-1,2,5,6-tetrahydropyridinoquinoline derivative," 24th ACS National Meeting, August 18-22, 2002, Boston, MA. Poster #258.					
	LC	NAKAZATO et al., "Design, synthesis and structure-affinity relationships of 4-methylidenepiperidine and 4-aryl-1,2,3,6-tetrahydropyridine derivatives as corticotropin-releasing factori receptor antagonists," <i>Bioorganic & Medicinal Chemistry</i> (2000) 8(5):1183-1193.					
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	LE	NICEWONGER et al., "Microwave-assisted acylation of 7-amino-5-aryl-6-cyanopyrido[2,3-d]pyrimidines," Molecular Diversity (2003) 7(2-4):247-252.					
	LF	NORMAN et al., "Structure –activity relationships of a series of pyrrolo(3,2-d) pyrimidine derivatives and related compounds as neuropeptide Y5 receptor antagonists" J. Med. Chem. (2000) 43(22):4288-4312.					
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	LK	PEDERSON, "The impact of obesity on the pathogenesis of non-insulin-dependent diabetes mellitus: a review of current hypotheses," <i>Diab. Metab. Rev.</i> , (1989) 5(6):495-509.					
Examiner Signature		Date Considered					

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	Application Number	10/541,657		
INFORMATION DISCLOSURE	Filing Date	January 14, 2004		
STATEMENT BY APPLICANT	First Named Inventor	Robert M. Jones		
	Art Unit	1624		
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	LL	PERRY et al., "Prospective study of risk factors for development of non-insulin dependent diabetes in middle aged British men," <i>BMJ</i> (1995) 310(6979):560-4.	
	LM	PHILLIPS et al., "Discovery of N-[2-[5-[Amino(imino)methyl]-2-hydroxyphenoxyl]-3,5-difluoro-6-[3-(4,5-dihydro-1-methyl-1H-imidazol-2-yl)phenoxy]pyridine-4-yl]-N-methylglycine(ZK-807834): A Potent, Selective, and Orally Active Inhibitor of the Blood Coagulation Enzyme Factor Xa," J. Med. Chem. (1998) 41(19):3557-3562.	
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	LO	POTENZA et al., "A rapid quantitative bioassay for evaluating the effects of ligands upon receptors that modulate cAMP levels in a melanophore cell line," <i>Pigment Cell Res.</i> (1992) 5(6):372-8.	
	LP	PRASAD, et al., "Convenient Methods for the Reduction of Amides, Nitriles, Carboxylic Esters, Acids and Hydroboration of Alkenes Using NaBH4/I2System," Tetrahedron (1992) 48(22):4623-4628.	
	LQ	PRESS et al., "Synthesis and SAR of 6-Substituted Purine Derivatives as Novel Selective Positive Inotropes," J. Med. Chem (1992) 35(24):4509-4515.	
	LR	QUINTELA et al., "6-Dimethylamino 1H-Pyrazolo[3,4-d]pyrimidine Derivatives as New Inhibitors of Inflammatory Mediators in Intact Cells," <i>Bioorganic & Medicinal Chemistry</i> (2003) 11:863-868.	
	LS	QUINTELA et al., "Pyrazolopyrimidines: synthesis, effect on histamine release from rat peritoneal mast cells and cytotoxic activity," Eur. J. Med. Chem. (2001) 36:321-332.	
	LT	RAM et al., "Chemotherapeutic agents. Part XXII. Synthesis of π -deficient pyrimidines as leishmanicides," <i>Indian Journal of Chemistry, Section B</i> (1991) 30B(10):962-965.	
	LU	REED et al., "In-vivo and in-vitro models of type 2 diabetes in pharmaceutical drug discovery," Diabetes Obes Metab, (1999) 1(2):75-86.	
	LV	REHWALD et al., "Syntheses of thieno[2,3-d]pyrimidines and aminopyrimidines from 2-alkoxy-5-cyano-4-thioxopyrimidine intermediates," <i>Heterocycles</i> (1998) 48(6):1157-1167.	
	LW	Remington's Pharmaceutical Sciences, 17th Ed., (1985), Mack Publishing Company, Easton, PA, p. 1418-1419.	

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Complete if Known Substitute for form 1449B/PTO January 14, 2004 **Application Number** INFORMATION DISCLOSURE July 6, 2006 Filing Date STATEMENT BY APPLICANT First Named Inventor Robert M. Jones Art Unit 1624 (Use as many sheets as necessary) To Be Determined Examiner Name of 22

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Sheet	18		of	22		Attorn	ey Docket Number	ARE	N34.US5.PCT	
					NON PATEN	T LITERAT	URE DOCUMENTS			
Examiner Initials *	Cite No. ¹		Include name of the author (in CAPITAL LETTERS), title of the article (when appropriate), title of the item (book, magazine, journal, serial, symposium, catalog, etc.), date, page(s), volume-issue number(s), publisher, city and/or country where published.						T²	
	*LX	Remir	igton's	Pharmace	eutical Sciences,	16 th Ed., (1	1980), Mack Publis	hing Com	pany, Easton, PA.	
	LY	Brom	EWCASTLE, et al., "Tyrosine Kinase Inhibitors. 10. Isomeric 4-[(3-romophenyl)amino]pyrido[d]pyrimidines are Potent ATP Binding Site Inhibitors of the Tyrosine Sinase Function of the Epidermal Growth Factor Receptor," J. Med. Chem. (1996) 39:1823-1835.							
	LZ	RAFF (1996)		ıl., "Diab	etes Mellitus,"	Principles	And Practice Of Mo	edical Gene	tics, 3 rd Ed. 1:1401-1440	
	MA	chlore	otrifluc	oro-, and			N-disubstituted ar pro-pyridines," Jou		luoro-, amino-3- Chemical Society [Section]	
	МВ	N,N-	ROBERTS et al., "Polychloroaromatic compounds. I. Oxidation of pentachloropyridine and its N,N-disubstituted amino derivatives with peroxyacids," Journal of the Chemical Society [Section] C: Organic (1968) (12):1537-1541.							
	МС	ROBI	NS, et		ential Purine Ar	ntagonists	. IV. Synthesis of S	Some 9-Me	thyl-6-substituted-	
	MD						clobutylamino de nii Nauk (1981) 34(1		f some aryl-substituted 5- 680.	
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	MF						nking region of the N Engl J Med. (198		sulin gene: a genetic 5-71.	
	MG		WELL (etrahydropyrid	yloxadiaz	oles: semirigid mu	iscarinic li	gands," J Med Chem (1991)	
	МН		AR et a		e and Efficient	Synthesis	of 6-(Hydroxymet	thyl)purine	es," Org. Lett. (2004)	
	MI SMITH et al., "Effects of positive allosteric modulators of the GABAB receptor on cocaine self-administration in rats," Psychopharmacology (2004) 173(1-2):105-111.					ptor on cocaine self-				
	MJ		SILVESTRI et al., "Novel indolyl aryl sulfones active against HIV-1 carrying NNRTI resistance mutations: synthesis and SAR studies," <i>J Med Chem</i> (2003) 46(12):2482-2493.							
	MK	1	NSMA 81-2283		A novel method	I for the sy	nthesis of aryl sul	fones," Te	trahedron Ltrs (2001)	
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Substitute	for form 1449E	B/PTO		Complete if Known			
WIE O		DIG	OLOGUEE	Application Number	10/541,657		
			CLOSURE	Filing Date	January 14, 2004		
STAT	EMENT	BYA	PPLICANT	First Named Inventor	Robert M. Jones		
				Art Unit	1624		
	(Use as man	y sheets as	necessary)	Examiner Name	To Be Determined		
Sheet	19	of	22	Attorney Docket Number	AREN34.US5.PCT		

		NON PATENT LITERATURE DOCUMENTS	
Examiner Initials *	Cite No. ¹	Include name of the author (in CAPITAL LETTERS), title of the article (when appropriate), title of the item (book, magazine, journal, serial, symposium, catalog, etc.), date, page(s), volume-issue number(s), publisher, city and/or country where published.	Τ2
	ML	STERNFELD et al., "Synthesis and serotonergic activity of 3-[2-(pyrrolidin-1-yl)ethyl]indoles: potent agonists for the h5-HT1D receptor with high selectivity over the h5-HT1B receptor," J Med Chem (1999) 42(4):677-690.	
	MM	STRUPCZEWSKI et al., "Synthesis and neuroleptic activity of 3-(1-substituted-4-piperidinyl)-1,2-benzisoxazoles," J Med Chem (1985) 28(6):761-769.	
	MN	SUAMI et al., "Nucleoside analogs. I. Synthesis of 1,3-dihydroxy-2-(6-substituted-9-purinyl)cyclohexane," Journal of Heterocyclic Chemistry (1969) 6(5):663-665.	
	МО	SUGIMOTO et al., "Preparation of Nitrogen-Containing π —Deficient Heteroaromatic Grignard Reagents: Oxidative Magnesiation of Nitrogen-Containing π -Deficient Halgenoheteroaromatics Using Active Magnesium," <i>J. Org. Chem.</i> (2003) 68:2054-2057.	
	MP	SUGIMOTO et al., "Lithiation of 1H-Pyrazolo[3,4-d]pyrimidine Derivative Using Lithium Alkanetellurolate," <i>Tetrahedron Letters</i> (1999) 40:2139-2140.	
	MQ	TERASHIMA et al., "Inhibition of human O6-alkylguanine-DNA alkyltransferase and potentiation of the cytotoxicity of chloroethylnitrosourea by 4(6)-(benzyloxy)-2,6(4)-diamino-5-(nitro or nitroso)pyrimidine derivatives and analogues," J Med Chem (1998) 41(4):503-508.	
	MR	THOMPSON et al., "N ⁶ ,9-Disubstituted Adenines: Potent, Selective Antagonists at the A1 Adenosine Receptor," <i>J. Med. Chem.</i> (1991) 34:2877-2882.	
	MS	THOMPSON et al., "Synthesis and evaluation of 6-(dibromomethyl)-5-nitropyrimidines as potential antitumor agents," J Med Chem (1997) 40(5):766-770.	
	МТ	TURCK et al., "Advances in the directed metallation of azines and diazines (pyridines, pyrimidines, pyrazines, pyridazines, quinolines, benzodiazines and carbolines). Part 2: Metallation of pyrimidines, pyrazines, pyridazines and benzodiazines," <i>Tetrahedron</i> (2001) 57(21):4489-4505.	
	MU	URGAONKAR et al., "Pd/P(i-BuNCH2CH2)3N: an efficient catalyst for Suzuki cross-coupling of aryl bromides and chlorides with arylboronic acids," <i>Tetrahedron Letters</i> (2002) 43(49):8921-8924.	
	MV	URWYLER et al., "N,N" –Dicyclopentyl-2-methylsulfanyl-5-nitro-pyrimidine-4,6-diamine (GS39783) and structurally related compounds: Novel allosteric enhancers of γ-aminobutyric acidB receptor function," Journal of Pharmacology and Experimental Therapeutics (2003) 307(1):322-330.	

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		211 DI 0		Application Number	10/541,657		
			CLOSURE	Filing Date	January 14, 2004		
ST	ATEMENT	T BY A	PPLICANT	First Named Inventor	Robert M. Jones		
				Art Unit	1624		
	(Use as man	y sheets as	necessary)	Examiner Name	To Be Determined		
She	et 20	of	22	Attorney Docket Number	AREN34.US5.PCT		

		NON PATENT LITERATURE DOCUMENTS	
Examiner Initials *	Cite No. ¹	Include name of the author (in CAPITAL LETTERS), title of the article (when appropriate), title of the item (book, magazine, journal, serial, symposium, catalog, etc.), date, page(s), volume-issue number(s), publisher, city and/or country where published.	T ²
	MW	VAUGHAN et al., "The Reformatsky Reaction. I. Zinc and Ethyl Alpha-Bromoisobutyrate," Dept. of Chem., The Univ. of Michigan, Ann Arbor, MI., (1964) 30:1790-1795.	
	MX	VICE, et al., "Concise Formation of 4-Benzyl Piperidines and Related Derivatives Using a Suzuki Protocol," J. Org. Chem. (2001) 66:2487-2492.	
	MY	VICE, et al., "Concise Formation of 4-Benzyl Piperidines and Related Derivatives Using a Suzuki Protocol," J. Org. Chem. (2001) 66:2487-2492, Supporting Information, pp. S1-S32	
	MZ	WANG et al., "Improving the oral efficacy of CNS drug candidates: discovery of highly orally efficacious piperidinyl piperidine M2 muscarinic receptor antagonists," J Med Chem (2002) 45(25):5415-5418.	
	NA	WELLS et al., "Regioselective nucleophilic substitutions of fluorobenzene derivatives," <i>Tetrahedron Letters</i> (1996) 37(36):6439-6442.	
	NB	WERBEL et al., "Synthesis and antimalarial effects of 5,6-dichioro-2-[(4-[[[4—(diethylamino) 1-methylbutyl]amino [[-6-methyl-2-pyrimidinyl)amino] benzimidazole and related benzimidazoles and I,H-Imidazo[4,5-b] pyridines," J. Het. Chem (1973) Vol. 10, 363-382.	
	NC	WILSON et al., "Microwave-assisted synthesis of 2-aminoquinolines," Tetrahedron Letters (2002) 43(4):581-583.	
	ND	WOLFE et al., "Scope and limitations of the Pd/BINAP-catalyzed amination of aryl bromides," <i>J</i> Org Chem (2000) 65(4):1144-1157.	
	NE	WOLFE et al., "Simple, efficient catalyst system for the palladium-catalyzed amination of aryl chlorides, bromides, and triflates," J Org Chem (2000) 65(4):1158-1174.	
	NF	WOLTER et al., "Copper-Catalyzed Coupling of Aryl Iodides with Aliphatic Alcohols," Organic Letters (2002) 4(6):973–976.	
	NG	WOLTER et al., "Copper-Catalyzed Coupling of Aryl Iodides with Aliphatic Alcohols," Organic Letters (2002) 4(6):973–976, Supporting Information, pp. S1-S16.	

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			CLOSURE	Filing Date	January 14, 2004	
STA	STATEMENT BY APPLICANT			First Named Inventor	Robert M. Jones	
				Art Unit	1624	
	(Use as man	y sheets as	necessary)	Examiner Name	To Be Determined	
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	NH	WU et al., "One-Pot Two-Step Microwave-Assisted Reaction in Constructing 4,5-Disubstituted Pyrazolopyrimidines," Org. Lett., (2003) 5(20):3587-3590.	
	NY	YAROVENKO et al., "New method for the preparation of 5-amino-1,2,4-oxadiazoles," Bull Acad Sci, USSR Div Chem Sci, (1991) 40:1924.	
	NZ	YOON et al., "Reaction of Diisobutylaluminum Hydride with Selected Organic Compounds Containing Representative Functional Groups," J. Org. Chem. (1985) 50:2443-2450.	
	OA	ZAMPONI et al., "Unique structure-activity relationship for 4-isoxazolyl-1,4-dihydropyridines," J Med Chem (2003) 46:87-96.	
	ОВ	ZAMPONI et al., "Unique structure-activity relationship for 4-isoxazolyl-1,4-dihydropyridines," J Med Chem (2003), Supporting Information., pp. 1-31.	
	ос	ZHANG, et al., "Preparation of 1-(Tri-n-Butylstannyl) Furanoid Glycals and Their Use in Palladium-Mediated Coupling Reactions," <i>Tetrahedron Letters</i> (1993) 34(10):1571-1574.	
	OD	ZHU et al., "Synthesis and mode of action of (125)I- and (3)H-labeled thieno[2,3-c]pyridine antagonists of cell adhesion molecule expression, J Org Chem. (2002) 67(3):943-8.	
	OE	Accession No. 2003:2415108 CHEMCATS, Interbioscreen Compound Library, Chemical Name: 1H-Pyrazolo[3,4-d]pyrimidine-4-amine, N-cyclohexyl-N-methyl-1-(3-methylphenyl)-, XP-002311326, 2003, CAS Registry No. 393844-90-1.	
	OF	Accession No. 2003:2415906 CHEMCATS, Interbioscreen Compound Library, Chemical Name: 1H-Pyrazolo[3,4-d]pyrimidine-4-amine, N-cyclohexyl-1-1-(4-methylphenyl)-, XP-002311325, 2003, CAS Registry No. 393844-89-8.	
	OG	Accession No. 2003:2416398 CHEMCATS, Interbioscreen Compound Library, Chemical Name: 1H-Pyrazolo[3,4-d]pyrimidine-4-amine, N-cyclohexyl-1-1-(2,4-dimethylphenyl)-N-methyl-, XP-002311324, 2003, CAS Registry No. 393844-91-2.	
	ОН	Accession No. 2003:2417080 CHEMCATS, Interbioscreen Compound Library, Chemical Name: 1H-Pyrazolo[3,4-d]pyrimidine-4-amine, N-cyclohexyl-N-methyl-1-phenyl)-, XP-002311323, 2003, CAS Registry No. 393844-87-6.	

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			SCLOSURE	Filing Date	January 14, 2004	
STA	STATEMENT BY APPLICANT			First Named Inventor	Robert M. Jones	
				Art Unit	1624	•
	(Use as mai	ny sheets as	s necessary)	Examiner Name	To Be Determined	
Sheet	22	of	22	Attorney Docket Number	AREN34.US5.PCT	

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	OI	Cover Sheet and 54 Compounds – CAS Registry file (23 pp.)	
	OJ	Cover Sheet and 18 Compounds – CAS Registry file (9 pp.)	
	OK	Cover Sheet and 2534 Compounds – CAS Registry and ChemCats files (817pp.)	
	OL	Cover Sheet and 1185 Compounds – CAS Registry and ChemCats Files (391pp.)	
	ОМ	23 Compounds - ChemCats File (11pp.)	
	*PI	Greene et al., Protective Groups in Organic Synthesis, 3rd Ed., John Wiley & Sons, New York (1999).	
 	*PJ	Remington, The Science and Practice of Pharmacy, 20th Ed., Lippincott Williams & Wilkins (2000).	
	*PK	Oae, Organic Chemistry of Sulfur, Ed., Plenum Press: New York (1977).	

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Signature	Considered	

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